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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO
10/773,472	02/05/2004	Daniella Licht	2609/68585-A/JPW/GJG/JBC 7030	
John P. White	7590 09/28/2007		EXAMI	NER
Cooper & Dunham LLP 1185 Avenue of the Americas New York, NY 10036			CHANNAVAJJALA, LAKSHMI SARADA	
			ART UNIT	PAPER NUMBER
			1615	
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Office Action Summary		Application No.	Applicant(s)				
		10/773,472	LICHT ET AL.				
		Examiner	Art Unit				
		Lakshmi S. Channavajjala	1615				
The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply							
WHIC - Exter after - If NO - Failu Any r	ORTENED STATUTORY PERIOD FOR REPLY CHEVER IS LONGER, FROM THE MAILING DANSIONS of time may be available under the provisions of 37 CFR 1.13 SIX (6) MONTHS from the mailing date of this communication. It period for reply is specified above, the maximum statutory period we re to reply within the set or extended period for reply will, by statute, reply received by the Office later than three months after the mailing and patent term adjustment. See 37 CFR 1.704(b).	ATE OF THIS COMMUNICATION 16(a). In no event, however, may a reply be tim rill apply and will expire SIX (6) MONTHS from cause the application to become ABANDONEI	N. nely filed the mailing date of this communication. D (35 U.S.C. § 133).				
Status							
·		action is non-final. ace except for formal matters, pro					
Dispositi	on of Claims						
5) 6) 7) 8)	Claim(s) 1-29,38-76,82-91,103-107 and 120 is/4a) Of the above claim(s) is/are withdraw Claim(s) is/are allowed. Claim(s) 1-29,38-76,82-91,103-107 and 120 is/Claim(s) is/are objected to. Claim(s) are subject to restriction and/or ion Papers	vn from consideration. /are rejected.					
10)□	The specification is objected to by the Examiner The drawing(s) filed on is/are: a) access Applicant may not request that any objection to the conference of Replacement drawing sheet(s) including the correction The oath or declaration is objected to by the Example 35 U.S.C. § 119	epted or b) objected to by the Edrawing(s) be held in abeyance. See non is required if the drawing(s) is obj	e 37 CFR 1.85(a). jected to. See 37 CFR 1.121(d).				
 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 							
2) Notice	t(s) e of References Cited (PTO-892) e of Draftsperson's Patent Drawing Review (PTO-948) mation Disclosure Statement(s) (PTO/SB/08) r No(s)/Mail Date <u>6-24-04, 7-2-04</u> .	4) Interview Summary Paper No(s)/Mail Da 5) Notice of Informal P 6) Other:	ate				

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DETAILED ACTION

Receipt of PRELIMINARY AMENDMENT dated 2-5-04, IDS dated 6-24-04 and 7-2-04 is acknowledged.

Claims 1-29, 38-76, 82-91,103-107 and 120 are pending.

Double Patenting

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. A nonstatutory obviousness-type double patenting rejection is appropriate where the conflicting claims are not identical, but at least one examined application claim is not patentably distinct from the reference claim(s) because the examined application claim is either anticipated by, or would have been obvious over, the reference claim(s). See, e.g., *In re Berg*, 140 F.3d 1428, 46 USPQ2d 1226 (Fed. Cir. 1998); *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970); and *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) or 1.321(d) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the conflicting application or patent either is shown to be commonly owned with this application, or claims an invention made as a result of activities undertaken within the scope of a joint research agreement.

Effective January 1, 1994, a registered attorney or agent of record may sign a terminal disclaimer. A terminal disclaimer signed by the assignee must fully comply with 37 CFR 3.73(b).

Claims 1-29, 38-76, 82-91,103-107 and 120 are provisionally rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 1-55 of copending Application No. 10/772,911 in view of US 4,704,285 ('285). Instant claims as well as the co-pending claims are directed to compressible tablets comprising valproic acid or its salts as an active agent, excipients such as HPMC,

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magnesium stearate, fillers, lubricants etc. Both sets of claims are directed to treating epilepsy, pain and conditions such as bipolar disorder with the above composition.

Instant claims differ from the co-pending claims in that instant claims HPMC that is different from the binder and also fail to claim the specific viscosity or the percentages of the methoxy or hydroxypropyl content of HPMC.

'285 teach compressible tablet preparation with HPMC ether fine particles as a matrix (col. 2, L 7-25). In addition to the active agents and HPMC ether, '285 teach inclusion of HPMC as a hydrocolloid and suggests that the particle size of HPMC is such that at least 70% pass through 100 mesh, with a HP content of 4%-12% and methoxy content of 19% to 30% (see col. 3, L 37-55). Thus, it would have been obvious for one of an ordinary skill in the art at the time of the instant invention was made to include HPMC such as that described by 285 in the compressible composition of instant claims because all the references are directed to compressible tablets and '285 teach that the claimed HPMC are routinely employed in compressible tablet preparation for improving the flow properties of the tablet. '285 further teach that the viscosity of HPMC is between 100 to 10,000 cps. Thus, the % of the MC and HP contents, viscosity and the particle sizes of instant claims are encompassed by the ranges of prior art. With respect to the claimed release rates, both sets of claims are directed to the same method of treatment with the same active compound and accordingly, optimizing the individual amounts of the components of the composition so as to achieve the desired release for an effective treatment with the active agent would have been obvious for a skilled artisan. This is a provisional obviousness-type double patenting rejection.

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Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

Claims 1-29, 38-76, 82-91,103-107 and 120 are rejected under 35 U.S.C. 103(a) as being unpatentable over US2001/0005512 to Anderson in view of Remingtons' Pharmaceutical Sciences (1990) and US 4,704, 285 OR US 6,419,953 to Qiu et al and Remingtons' Pharmaceutical Sciences (1990) and US 4,704, 285 ('285).

Anderson teaches a pharmaceutical composition comprising valproate compounds such as divalproex sodium as an active agent. For the dosage forms containing the active agent, Anderson teaches tablet formulations comprising the active agent and hydroxypropyl cellulose, which read on the instant components I) and ii). For the instant filler, Anderson teaches microcrystalline cellulose and lactose in the above

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dosage form. For the instant lubricant, Anderson teaches magnesium stearate (paragraphs 0107 – 0114). Anderson also teaches the active agents for the same treatment i.e., epilepsy and bipolar disorder (col. 1-2). While instant claims recite an immediate release composition in the preamble, the claims do not distinguish or specify as to what the release rate is or how quickly and how much of the active agent is released. Accordingly, the preamble does not add any patentable weight to the claimed composition. Further, Anderson teaches the various tabletting ingredients as percentages of the total weight of the tablet as opposed to the amounts. However, the tablets of Anderson are prepared in the same manner (compression tablets) as that claimed in the instant i.e., admixing the predetermined amounts and compressing the tablets. Anderson also teaches the excipients for the same purpose i.e., filler, lubricant etc and accordingly, optimizing the amount of an excipient with an expectation the desired tabletting effect such as lubrication, increasing the bulk (with a filler) etc., would have been within the scope of a skilled artisan.

Qiu et al (Qiu) teaches controlled release composition comprising an antiepileptic agent, valproic acid or its salts such as an ester, amide etc., prepared by intimately mixing the components of the composition and compression method (lines bridging col. 2-3 & col. 5, L 8-20). The composition, in the form of tablets, contains hydroxypropyl methylcellulose (examples formulations) and excipients such as magnesium stearate, lactose, microcrystalline cellulose (col. 3, L 1-53 & col. 5).

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Both Anderson and Qiu fail to teach the claimed disintegrant and also the specific HPMC with the claimed percentages of HP and MC contents, viscosity, particle sizes etc., in the composition comprising valproic acid or its salts. However, both the references are directed to preparing compressed dosage forms for a controlled release of active agent.

Remingtons' Pharmaceutical Sciences (Remingtons') teach oral dosage forms, particularly, compressed tablets comprising the tabletting excipients such as diluents, binders, disintegrants, glidants etc (pages 134-1637). Remingtons' teach that a substance or a mixture of substances is added to a tablet so as to facilitate its break up or disintegration after administration (page 1637). Among the disintegrants, Remingtons' teach the instant claims sodium starch glycolate or croscarmellose sodium as super Disintegrants, which are also claimed in the instant application (page 1637). Further, Remingtons' suggests that the disintegrant may be added to the active ingredients along with the lubricants, fillers etc., or may be divided into two portions, in which one portion is added prior to granulation and the other is added and the reminder mixed with lubricants before compression.

Further, '285 teach compressible tablet preparation with HPMC ether fine particles as a matrix (col. 2, L 7-25). In addition to the active agents and HPMC ether, '285 teach inclusion of HPMC as a hydrocolloid and suggests that the particle size of HPMC is such that at least 70% pass through 100 mesh, with a HP content of 4%-12% and methoxy content of 19% to 30% (see col. 3, L 37-55).

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Thus, it would have been obvious for one of an ordinary skill in the art at the time of the instant invention to use a disintegrant such as sodium starch glycolate or croscarmellose sodium, of Remingtons' in the compression tabletting composition of Anderson or Qiu because Remingtons' teach that the disintegrants or super disintegrants allow the guick break up of the tablet for its rapid dissolution and super disintegrants such as croscarmellose sodium are even more effective because of their activity even at low amounts and high swelling. Accordingly, depending on the desired rapidity of drug release, a skilled artisan would have employed an appropriate amount of a disintegrant in the composition of Anderson or Qiu. Further, with respect to the composition claims containing specific amounts of fillers, active agent and the disintegrants, absent any unexpected result optimizing the amounts of each of the active agent or the tabletting ingredients (lubricant, filler, disintegrant, release polymer) so as to achieve the desired release rate would have been within the scope of a skilled artisan. With respect to the claimed HPMC viscosity, % of HP and MC and particles, it would have been obvious for one of an ordinary skill in the art at the time of the instant invention was made to include HPMC such as that described by 285 in the compressible composition of Anderson or Qiu because all the references are directed to compressible tablets and '285 teach that the claimed HPMC are routinely employed in compressible tablet preparation for improving the flow properties of the tablet. '285 further teach that the viscosity of HPMC is between 100 to 10,000 cps. Thus, the % of the MC and HP contents, viscosity and the particle sizes of instant claims are encompassed by the ranges of prior art. For the specific release rates claimed, both

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Anderson and Qiu recognize the importance of valproic acid in treating the claimed conditions such as epilepsy and also teach the claimed excipients. Accordingly, optimizing the individual amounts of the components of the composition so as to achieve the desired release for an effective treatment with the active agent would have been obvious for a skilled artisan.

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Any inquiry concerning this communication or earlier communications from the examiner should be directed to Lakshmi S. Channavaijala whose telephone number is 571-272-0591. The examiner can normally be reached on 7.00 AM -4.00 PM.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Michael Woodward can be reached on 571-272-8373. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

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September 22, 2007

PRIMARY EXAMINER